# An Immunoglobulin Superfamily-Like Domain Unique to the *Yersinia pseudotuberculosis* Invasin Protein Is Required for Stimulation of Bacterial Uptake via Integrin Receptors

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The binding of the Yersinia pseudotuberculosis and Yersinia enterocolitica invasin proteins to  $\beta_1$  integrin receptors allows internalization of these organisms by cultured cells. The C-terminal 192-residue superdomain of the Y. pseudotuberculosis invasin is necessary and sufficient for integrin recognition, while a region located outside, and N-terminal to, this superdomain strongly enhances the efficiency of bacterial uptake. Within the enhancer region is a domain called D2 that allows invasin-invasin interaction. To investigate the role of the enhancer region, bacterial cell binding and entry mediated by the Y. pseudotuberculosis invasin protein (invasin<sub>pstb</sub>) was compared to that of Y. enterocolitica invasin (invasin<sub>ent</sub>), which lacks the D2 self-association domain. Invasin<sub>ent</sub> was shown to be unable to promote self-interaction, using the DNA binding domain of  $\lambda$  repressor as a reporter. Furthermore, two genetically engineered in-frame deletion mutations that removed D2 from invasin<sub>pstb</sub> were significantly less proficient than wild-type invasin<sub>pstb</sub> at promoting uptake, although the amount of surface-exposed invasin as well as the cell binding capacity of the recombinant Escherichia coli strains remained similar. Competitive uptake assays showed that E. coli cells expressing invasin<sub>pstb</sub> had a significant advantage in the internalization process versus either E. coli cells expressing invasin<sub>ent</sub> or the invasin<sub>pstb</sub> derivatives deleted for D2, further demonstrating the importance of invasin self-interaction for the efficiency of invasin-mediated uptake.

Many bacterial pathogens have adopted strategies to adhere to and efficiently penetrate normally nonphagocytic host cells (3, 8, 15). Entry into host cells permits bacteria to either grow and multiply in a protected niche (30) or to gain access to other tissues within the host (11, 30). The latter tactic is used by a number of enteric pathogens that translocate into subepithelial sites, allowing the initiation of systemic disease. A favored cell type for the spread of pathogens from the intestine is the M cell, which can be found interdigitated within the epithelium overlying lymphoid Peyer's patches in the small intestine (31). In the case of enteropathogenic Yersinia organisms, the tropism for this cell type can partly be explained by the fact that no other cell in the intestinal epithelium efficiently presents the receptors that recognize the bacterial invasin protein, which is required for efficient translocation into Peyer's patches (25, 31, 33).

Invasin is encoded by both *Yersinia enterocolitica* and *Yersinia pseudotuberculosis* and apparently plays a similar role in both organisms, allowing bacterial colonization of regional lymph nodes after ingestion (25, 33). The protein is localized in the outer membrane, with the C-terminal 479 amino acids of the *Y. pseudotuberculosis* invasin protein (invasin<sub>psth</sub>) exposed on the bacterial cell surface (22, 23). This region is responsible for promoting cell adhesion and uptake by binding to multiple  $\beta$ -chain integrin receptors on the host eukaryotic cell (16). The N-terminal half of invasin appears to be required for export of

the hydrophilic C-terminal region across the outer membrane (22).

The integrin receptor family consists of several related heterodimeric integral membrane proteins, involved in various adhesive functions, including cell-cell interaction, cell migration, cellular differentiation, and attachment to extracellular matrix proteins (14). Integrin receptors, via their cytoplasmic domains, are capable of signaling to cytoskeletal components after adhering to substrates. Invasin-mediated bacterial uptake is inhibited by drugs that antagonize either actin polymerization or tyrosine phosphorylation (9, 36). One tyrosine-phosphorylated mammalian protein that is clearly required for uptake is FAK (1). In addition, determinants within the cytoplasmic domain of the integrin that allow association of the receptor with the cytoskeleton and endocytic components modulate the efficiency of bacterial uptake (41).

The crystal structure of the C terminus of Y. pseudotuberculosis invasin has been determined, extending from residues 503 to 986 (12). The protein is arrayed as a series of five domains, extending in a rod-like 180-Å structure (see Fig. 1A). The first four domains (D1 to D4) are predominantly β stranded, each adopting a folding topology found in members of the immunoglobulin superfamily. The fifth domain (D5), which has interspersed  $\alpha$ -helical and  $\beta$ -stranded regions, is related to Ctype lectin-like domains (42). The minimal region of invasin required for binding to integrin receptors contains just D4 and D5 (22). D4 and D5 have a large interdomain interface, resulting in the formation of a superdomain extending from residues 795 to 986. Residues from both D4 and D5 appear to be presented to the integrin receptor, because point mutations in both domains result in defective receptor binding (24, 37). A derivative containing just D4 and D5, when coated as a monomer on the surface of latex bead, is inefficient at promoting

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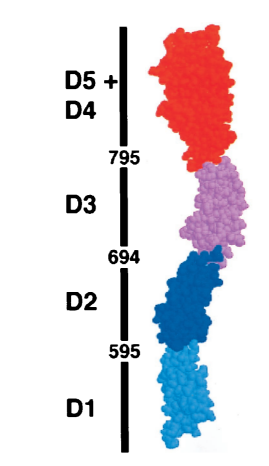


FIG. 1. (A) Spacefilling representation of the invasin $_{pstb}$  crystal structure. Different domains are indicated as D1 through D5 and depicted as different colors. Residue numbers that are located in interdomain regions are noted next to the structure. The region sufficient to promote cI multimerization is shown in dark blue and represents the entire D2 region in the crystal structure. The minimal region necessary for integrin recognition is shown in red and is the D4-D5 superdomain (12). (B) Comparison of a portion of Y. pseudotuberculosis invasin sequence and selected orthologs from other gram-negative bacteria. The predicted amino acid sequence (amino acids 555 to 734) of invasin<sub>pstb</sub> (19), extending from the middle of D1 to the middle of D3, is shown in panel A. This is aligned with the sequences of Y. enterocolitica invasin (45) and the enterohemorrhagic E. coli (EHEC) intimin. The Citrobacter freundii and enteropathogenic E. coli intimins align similarly to the EHEC protein (data not shown). The amino acid sequences of two genetically engineered invasin $_{pstb}$  mutants cloned on plasmids pPD254 and pPD255 are shown below. Sequences were aligned by the program Clustal W (2). Residue numbers of each protein are given at the start and the end of each line. Identical amino acid residues found among all proteins are noted with asterisks (\*). Similar amino acid residues are enclosed by shaded boxes. The boxed sequence of the Y. pseudotuberculosis invasin represents the domain responsible for multimerization and is coincident with the endpoints of D2 in the crystal structure (12). The dashes indicate the regions that are deleted in invasinent and intimins.



uptake of integrin-bound particles (7). As beads coated by D1 to D5 can be internalized efficiently, residues within D1 and D3 enhance uptake.

An activity that may play a role in enhancing uptake has been identified in D2. Hybrid proteins containing only residues from this domain are capable of promoting homotypic interaction (7). No other portion of the determined structure is capable of this activity. That invasin self-interaction appears to play a role in uptake is supported by the fact that beads coated with D4 and D5 are internalized if this superdomain is dimerized by binding to immunoglobulin G-coated particles (7).

The ability of  $invasin_{pstb}$  D2 to self-associate and the importance of D1 to D3 in promoting efficient uptake has led to the proposal that direct interaction of multimeric invasin with mul-

tiple integrins mediates receptor clustering, resulting in an intracellular signal for bacterial uptake (7). The *Y. enterocolitica* invasin protein (invasin<sub>ent</sub>), which exhibits an overall homology of 85% to invasin<sub>pstb</sub>, has an internal deletion of 99 amino acids that completely removes D2 (Fig. 1B) (45). Several sequence comparison programs place a similar deletion within the intimins, bacterial cell attachment proteins that have high sequence similarity to invasin (10, 20, 39). The recently described global fold of the carboxyl-terminal 280-amino-acid fragment of one such intimin is highly similar to the folding pattern of invasin domains D3 to D5, emphasizing that these proteins are part of a single family (21, 26). Even so, intimins are able to mediate bacterial adhesion to host cells in the absence of a homologous D2 region.

TABLE 1. Bacterial strains, plasmids, and bacteriophages

Strains, plasmids, and phage	Description <sup>a</sup>	Reference
Strains		
E. coli K12		
XL1blue	F'::Tn10 pro $A^+B^+$ lac $I^q$ $\Delta(lacZ)$ M15/recA1 endA1 gyrA96 (Nal <sup>r</sup> ) thi hsdR17 ( $r_K^ m_K^+$ )	6
	supE44 relA1 lac	
JH372	lysogenic $\lambda 202$ (imm 21 P <sub>B</sub> -lacZ $\Delta O_{B^2}$ )	13
SR2	$F^-$ araD $\Delta(lacU)$ $\Delta phoA$ lpp5508 galE rpsL galE galK degP::Tn5	35
5112	- map =(mee) =p.m. ppeede 8m2 .p. 2 8m2 8m1 mg1 .r. m	
Y. pseudotuberculosis	Type III(P <sup>+</sup> ), pIB1	4
Y. enterocolitica 8081 c		32
Plasmids		
pBR322	Cloning vector, Ap <sup>r</sup>	5
pHSG575/576	pSC101-based cloning vectors, placZ MCS, lacZ', Cm <sup>r</sup>	40
pMin3	Produces MBP-Inv <sub>ent</sub> fusion protein, Ap <sup>r</sup>	32
pFG157	λ cI ind1, lacUV5 promoter, colE1, ori, Ap <sup>r</sup>	13
pJH370	$\lambda$ cI ind1 aa 1–132 fused to zipper domain of GCN4, Ap <sup>r</sup>	13
pJH391	$\lambda$ cI ind1 aa 1–132 fused to lacZ, Ap <sup>r</sup>	13
pKH101	$\lambda cI indI$ aa 1–132 <sup>b</sup> Ap <sup>r</sup>	13
pPD207	pHSG576, inv <sub>pstb</sub> +, Ap <sup>r</sup>	This work
pPD208	Produces cI <sub>N</sub> -Inv509-986-Inv <sub>pstb</sub> 478, Ap <sup>r</sup>	7
pPD210	Produces cI <sub>N</sub> -Inv795-986-Inv <sub>pstb</sub> 202, <sup>b</sup> Ap <sup>r</sup>	7
pPD226	Produces cI <sub>N</sub> -Inv575-694, <sup>b</sup> Ap <sup>r</sup>	7
pPD229	Produces cI <sub>N</sub> -Inv <sub>ent</sub> 456-835, <sup>b</sup> Ap <sup>r</sup>	This work
pPD230	Produces $cI_{N}$ -Inv <sub>pstb</sub> 575-694-Inv <sub>em</sub> 456-835, <sup>b</sup> Ap <sup>r</sup> pHSG576 $inv_{ent}$ <sup>+</sup> , Cm <sup>r</sup>	This work
pPD231	pHSG576 inv <sub>ent</sub> <sup>+</sup> , Cm <sup>r</sup>	This work
pPD244	Produces cI <sub>N</sub> -Inv596-694-Inv <sub>ent</sub> 380, <sup>b</sup> Ap <sup>r</sup>	7
pPD254	$pPD207, inv_{pstb}(\Delta 1-595-697),^{c} Cm^{r}$	This work
pPD255	$pPD207, inv_{pstb}(\Delta 2-594-697),^{d} Cm^{r}$	This work
pPD256	pPD207, Ap <sup>r</sup> , Cm <sup>s</sup>	This work
pRI203	pBR325 $inv^+$ , Ap <sup>r</sup>	19
pRI253	$pT7-4 inv^+, Ap^r$	18
pRI285	Produces MBP-Inv497 <sub>pstb</sub> fusion protein	24
pZ150	Cloning vector carrying lacUV5 promoter, Apr	13
Dostonionkon		
Bacteriophage λKH54	$\Delta c \mathrm{I}$	13

a aa, amino acid; Apr, ampicillin resistance; Cmr, chloramphenicol resistance; Cms, chloramphenicol sensitivity; Nalr, nalidixic acid resistance.

In this study, the activities of invasin derivatives lacking D2 were examined. The lack of this domain in the wild-type invasin<sub>ent</sub> protein was found to lower the efficiency of bacterial internalization relative to invasin<sub>pstb</sub> and eliminate efficient homotypic interaction. This difference in the activity of the two proteins may contribute to the observed differences in the colonization pattern of these two organisms within intestinal lymph nodes, as *Y. enterocolitica* efficiently proliferates and forms large abscesses in these sites, whereas *Y. pseudotuberculosis* does not.

# MATERIALS AND METHODS

Bacterial strains, cell culture, and media. Bacterial strains, phages, and plasmids used in this study are described in Table 1. Bacterial strains were grown with aeration in Luria-Bertani broth (Difco) or in M9 medium containing 0.4% glucose (27, 38) at  $37^{\circ}$ C (Escherichia coli) or at  $28^{\circ}$ C (Yersinia). The antibiotics used for bacterial selection were as follows: ampicillin,  $100 \,\mu g/ml$ ; chloramphenicol,  $30 \,\mu g/ml$ ; tetracycline,  $5 \,\mu g/ml$ ; kanamycin,  $50 \,\mu g/ml$ ; gentamicin,  $50 \,\mu g/ml$ . HEp-2 cells were cultured in RPMI 1640 media (Irvine Scientific) supplemented with 5% newborn calf serum (Life Technology Inc.) and  $2 \, mM$  glutamine at  $37^{\circ}$ C in the presence of 5% CO<sub>2</sub>.

**Nucleic acid techniques.** Preparations of small-scale plasmid DNA, restriction digestions, ligations, and transformations were performed as previously described (27, 38). Large-scale plasmid preparations were purified on Qiagen columns (Hilden, Germany) according to manufacturer's protocols. PCR products were purified with the QIAquick Kit (Qiagen) before and after restriction digestions.

PCRs were performed in a 100- $\mu$ l mixture consisting of 1× Taq polymerase buffer, 0.2 mM concentration of deoxynucleoside triphosphate (Pharmacia Biotech), 10 pmol of each primer, 0.5  $\mu$ g of template DNA, 5 mM MgCl<sub>2</sub>, and 2.5 U of recombinant Taq polymerase (Perkin-Elmer Cetus). The reactions were performed as follows: denaturation at 95°C for 5 min, 20 cycles annealing at 58°C for 30 s, extension at 72°C for 30 s to 2 min (depending on the length of the expected PCR product), and 95°C for 30 s, in a DNA thermal cycler PTC-200 (MJ Research).

Plasmids and oligonucleotides. Plasmids used in this study are listed in Table 1. Plasmid pPD207 ( $inv_{pstb}^+$ ) was derived by inserting the EcoRI-HindIII fragment of pRI253 (17), carrying the entire  $inv_{pstb}$  gene into the EcoRI and HindIII sites of pHSG576 (40). Plasmid pPD231 ( $inv_{ent}^+$ ) was constructed by inserting a PCR fragment harboring the  $inv_{ent}$  gene, including the  $inv_{ent}$  promoter, into vector pHSG575 that was cut with EcoRI and BamHI. This product was generated by using the upstream primer  $inv_{ent}$  (5'-CCATATGAATTCCTTAACTAA GCCAGCGGTTGC-3') (creates an EcoRI site) and downstream primer  $inv_{ent}$  (5'-AACGGTGGATCCCGGCAACCTGCATAACGGGC-3') (creates a BamHI site). The in-frame deletion derivatives of  $inv_{pstb}$  that mimic the inv sequence of Y. enterocolitica were constructed by using the Transformer site-

 $<sup>^</sup>b$  Protein expressed from the *lacUV5* promoter; the numbers of the range indicate the amino acids of *inv*<sub>pstb</sub> fused to cI<sub>N</sub>'.

<sup>&</sup>lt;sup>c</sup> The numbers indicate the amino acids of *inv<sub>pstb</sub>* deleted in the construct.

<sup>&</sup>lt;sup>d</sup> The numbers indicate the amino acids of  $inv_{pstb}$  deleted in the construct; in addition, amino acids AVLP from the equivalent Y. enterocolitica sequence were introduced into this region (see Fig. 1).

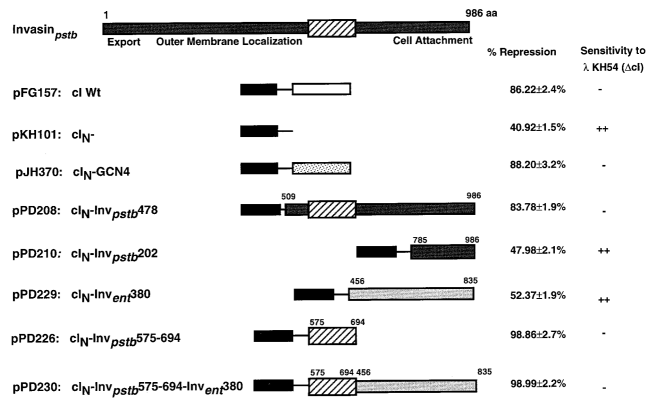


FIG. 2. Repressor activity of  $cI_N$ -Inv chimeric proteins. Schematic representations of invasin and the chimeric  $cI_N$ -Inv proteins are shown. The  $cI_N$  portion of the chimeric proteins (DNA binding domain) is shown as a black box, and the dimerization domain of  $\lambda$  repressor is shown as white box. The invasin sequence of Y pseudotuberculosis is represented by dark gray boxes, with the exception of the region encoding domain 2, which is cross-hatched. The sequences of Y enterocolitica are shown as light gray boxes, which are aligned in relationship to the invasin wild-type sequence. Repression activity of the chimera and sensitivity to phage KH54( $\Delta cI$ ) were measured as described (see Materials and Methods). The efficiency of KH54( $\Delta cI$ ) plating represents sensitivity to a phage lysate of  $10^9$  phages/ml. –, no plaque formation; +, plaque formation.

directed mutagenesis kit (Clontech). For selection, primer 5'-GGGAAAACTG TCCTGCAGCACAGATGAAAACGG-3' (creates a PstI site) was used, designed to change the unique NdeI site in pHSG576 into a PstI site. For pPD254 and pPD255, primers 5'-GTTGATACCGACTTTGTT-Δ-CCTATTCCAGATG CTGGC-3' and 5'-GTTGATACCGACTTT GCC GTG CTG CCG CCTATTC CAGATGCTGGC-3' were used as mutagenic primers (Δ labels the deletion introduced in inv<sub>pstb</sub>, underlined codons show the AVLP sequence introduced at the deletion site of inv<sub>pstb</sub> [Fig. 1B]). To change the antibiotic resistance of pPD207 and generate pPD256, the bla gene of pBR322 was amplified with primers 5'-GCGGCGCCATGGGGAAATGTGCGCGGAACC-3' and 5'-GCGGCCCATGGTTGACAGTTACCAATGC-3' (creating NcoI sites) and inserted into the unique NcoI site in the chloramphenicol resistance gene of pPD207.

To construct the chimeric cI<sub>N</sub>-Inv<sub>ent</sub>380 protein encoded by plasmid pPD229, the DNA-binding domain of cI (cI<sub>N</sub>) was fused to the C-terminal 380 amino acids of invasin<sub>ent</sub>. To this end, a fragment of the *inv* gene of *Y. enterocolitica* (*inv*<sub>ent</sub>) encompassing amino acids 456 to 835 was amplified by PCR using upstream primer 5'-TATTACGTCGACAGTGACTGATGATGGTGCGCTTGC-3' (creating a *Sal*I site) and downstream primer 5'-AACGGTGGATCCCGGCAACC TGCATAACGGGC-3' (creating a *Bam*HI site) and cloned into the *Sal*I and *Bam*HI site of pJH391 encoding cI<sub>N</sub> (13). To generate cI<sub>N</sub>-Inv<sub>pstb</sub>575-694-Inv<sub>ent</sub>380 encoded by plasmid pPD230, a PCR-derived fragment carrying invasin<sub>pstb</sub>, codons 575 to 694, was inserted into the *Sal*I site upstream of the *inv*<sub>ent</sub> gene on pPD229, using primers 5'-GGGGGGTCGACGGTGATAACCACCA GTAATGGTGCG-3' and 5'-GGGGGGGTCGACGGTGATCACCACCA GTAATGGTGCG-3' and 5'-GGGGGGGGTCGACGGATCTGCCGTGAAA TTAACCG3', both creating *Sal*I sites. A schematic representation of the cI<sub>N</sub>-inv constructs is shown in Fig. 2. Oligonucleotides were synthesized by the Howard Hughes Medical Institute Microchemical Facility (Harvard Medical School, Boston, Mass.).

Overexpression of invasin homologues. For cell binding and cellular uptake experiments with  $E.\ coli$  cells harboring invasin derivatives, bacteria were grown in M9 medium containing 0.4% glucose and antibiotic to an absorbance at 600 nm ( $A_{600}$ ) of 0.7. Induction of invasin expression was performed by adding 2 mM IPTG (isopropyl- $\beta$ -D-thiogalactopyranoside) (Sigma) for 60 min to the growth medium. Bacteria used in cell binding assays were adjusted to the identical

densities, and viable counts were performed on each culture to determine the exact multiplicities of infection.

Expression and purification of MBP-Inv fusion proteins. The purification of the MBP-Inv hybrid proteins were performed as previously described, with slight modifications (7). One liter of E. coli SR2 cells carrying either plasmid pMin3 or pRI285 was grown at 28°C in L broth to an  $A_{600}$  of 0.7 before adding IPTG to a final concentration of 2 mM. The cells were then grown at 28°C for 2 additional hours before being harvested, and all following procedures were performed at 4°C. Cells were resuspended in 5 ml of 10 mM Tris (pH 8.0) with a protease inhibitor cocktail containing 5 mM phenylmethylsulfonyl fluoride, 10 mM Pepstatin (Sigma), 10 mM E64 (Boehringer Mannheim), 20 µM Leupeptin (United States Biochemical), and 10 µM Chymostatin (Sigma). Subsequently, the cells were lysed by sonication (Branson Instruments; 50% pulse, 5 min). The soluble MBP-Inv protein extract was separated from insoluble cell material by centrifugation at  $25,000 \times g$  and purified by affinity chromatography on cross-linked amylose as described previously (22, 35). MBP-Inv497 was purified by ionexchange chromatography (7, 24). Protein concentrations were determined by the bicinchoninic acid protein assay (Pierce).

Preparation of total cell extracts, gel electrophoresis, and Western blotting. Cultures of E. coli XL1blue cells harboring the various inv alleles on plasmids pPD207, pPD231, pPD254, and pPD255 were grown overnight at 37°C in Luria-Bertani medium. The optical density was adjusted, and a 1-ml aliquot was withdrawn from each culture. The cells were collected by centrifugation, resuspended in 100 μl of sample buffer (0.06 M Tris [pH 6.8], 2% sodium dodecyl sulfate [SDS], 10% glycerol, 3% dithiothreitol, 0.001% bromphenol blue) and subsequently lysed by incubation at 100°C for 5 min. To reduce the viscosity of the total cell extracts, 3 µl of benzon nuclease (Merck, Darmstadt, Germany) was added to the samples and incubated at 37°C for 10 min. For the immunological detection of the invasin proteins, 2-µl portions of the total cellular extracts were loaded onto SDS-10% polyacrylamide gels, and the proteins were separated by electrophoresis and transferred to an Immobilon membrane (Millipore). The bound proteins were then probed with a monoclonal invasin<sub>pstb</sub>- or polyclonal invasin<sub>ent</sub>-specific antiserum. The antigen-antibody complexes were visualized with a second goat alkaline-phosphatase-conjugated antibody (Sigma) using

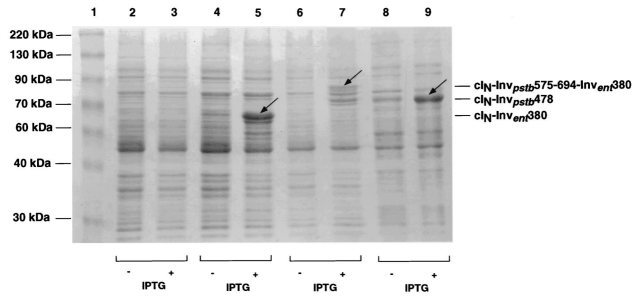


FIG. 3. Expression of the  $cI_{N}$ -inv gene products of Y. pseudotuberculosis and Y. enterocolitica in E. coli. Total cell extracts were prepared from strain JH372 carrying the  $cI_{N}$ -inv fusions grown either in the presence (+IPTG) or absence (-IPTG) of inducer. The proteins were electrophoretically separated on an SDS-12% polyacrylamide gel, and the gel was stained with Coomassie brilliant blue. The arrows indicate the position of the  $cI_{N}$ -Inv hybrid proteins and a molecular mass standard is shown in lane 1. Lanes 2 and 3, JH372 pKH101; lanes 4 and 5, JH372 pPD229 ( $cI_{N}$ -Inv $_{ent}$ 380); lanes 6 and 7, JH372 pPD230 ( $cI_{N}$ -Inv $_{pstb}$ 575-694-Inv $_{ent}$ 380); lanes 8 and 9, JH372 pPD208 ( $cI_{N}$ -Inv $_{pstb}$ 478).

5-bromo-4-chloro-3-indolylphosphate (XP) and nitroblue tetrazolium (Boehringer) as substrates.

ELISA to determine the invasin surface concentration. Enzyme-linked immunosorbent assays (ELISAs) were used to compare the surface concentration of invasin in the outer membranes of Y. pseudotuberculosis and Y. enterocolitica cells. Serial dilutions of  $2 \cdot 10^7$  bacteria were incubated in phosphate-buffered saline (PBS) containing 2% goat serum, 1 µg of primary anti-invasin antibodies (mAb3A2 for invasin<sub>nsth</sub> or a polyclonal invasin<sub>ent</sub> antiserum) per ml, followed by reprobing with 5 µg of goat anti-mouse or anti-rabbit immunoglobulin G alkaline phosphatase (Zymed) per ml for 1 h at room temperature. After washing, the cell density was redetermined at  $A_{600}$  and the bacteria were incubated in AP-buffer with 1 mg of  $\sigma^{104}$  alkaline phosphate substrate (Sigma) per ml in AP-buffer containing 100 mM Tris-HCl (pH 9.5), 5 mM MgCl<sub>2</sub>, 150 mM NaCl<sub>2</sub>. Exactly 20 min after starting the color reaction, the assay was monitored at 405 nm with a microtiter spectrophotometer (Bio-Rad). The amount of invasin on the bacterial cell surface was determined from standard ELISAs using the equivalent purified MBP-Invasin fusion protein. To do so, plastic wells were coated with serial dilutions of the purified fusion protein for 16 h at 4°C. The amount of bound protein was determined by subtracting the amount of protein that remained in the supernatant, determined by bicinchoninic acid protein assay. The relative amount of surface-exposed invasin concentration was determined by comparing the color reaction of the bacteria harboring invasin with that of the purified proteins, using concentrations that yielded a linear relationship between protein concentration and color reaction. Subsequently, the calculated invasin concentration was normalized to cell number. The corrected values shown (see Fig. 4C) are expressed relative to the amounts of surface-exposed invasin of uninduced bacteria XL1blue pPD207 (inv<sub>pstb</sub>+), which is defined as 1.0.

Cell binding and uptake assay. In preparation for cell binding or uptake assays,  $5 \times 10^4$  HEp-2 cells were seeded and grown overnight in individual wells of 24-well cell culture plates (Costar), using round coverslips when appropriate. Cell monolayers were washed three times with PBS and incubated in RPMI 1640 medium supplemented with 20 mM HEPES (pH 7.0) and 0.4% bovine serum albumin before the addition of approximately 5  $\times$  10  $^6$  bacteria. For the competitive invasion assays, equal amounts of  $2.5 \times 10^6$  bacteria were premixed before addition to cells. Bacteria were centrifuged onto the cell monolayer (1,000 rpm. 5 min in a Hermle tabletop centrifuge) and incubated at 20°C to test for cell binding (and prevent bacterial uptake) or at 37°C to test for invasion. To assay cell binding, 1 h postinfection, the cells were washed three times with PBS and fixed with 2% paraformaldehyde in PBS for 20 min at room temperature. Subsequently, the samples were washed three times with PBS and mounted in PBS containing 0.1% p-phenylenediamine and 80% glycerol (vol/vol). Quantification of adherent bacteria was performed by using a Zeiss Axioskope (Jena, Germany). The total number of bacteria associated with the eukaryotic cell was determined by counting bacteria bound per HEp-2 cell under phase-contrast microscopy or with Giemsa-stained preparations. Two hundred individual cells were counted, and the number of adherent bacteria per cell was determined as the mean of 50 individual cells. Bacterial uptake was assessed 90 min after infection as the percentage of bacteria which survived killing by the addition of the antibiotic gentamicin to the external medium, as described previously (7). For each strain, the relative level of bacterial uptake was determined by calculating the number of CFU that arose relative to the total number of bacteria introduced onto monolayers, performed in triplicate.

#### **RESULTS**

The presence of D2 in the *Y. pseudotuberculosis* invasin protein and its absence in the *Y. enterocolitica* invasin suggested that these two proteins may differ in their relative abilities to either promote cellular uptake or undergo homotypic interaction (Fig. 1) (45). To further elucidate the role of the *Y. pseudotuberculosis* invasin D2 region, the ability of the invasin of *Y. enterocolitica* to mediate homotypic interaction was determined and the ability of the two homologs to promote bacterial uptake was compared.

Y. enterocolitica invasin does not promote self-interaction in the one-hybrid assay. In previous experiments, the region responsible for  $invasin_{pstb}$ -invasin<sub>pstb</sub> interaction was mapped to amino acids 595 to 694, which is now known to define the endpoints of D2 (7, 13) (Fig. 1A, dark blue domain). To investigate whether the Y. enterocolitica invasin is able to promote self-interaction, the monomeric DNA-binding domain of cI (cI<sub>N</sub>) was fused to the C-terminal 380 amino acids of inva- $\sin_{ent}$  (cI<sub>N</sub>-Inv<sub>ent</sub>380) and tested in the  $\lambda$  repressor one-hybrid assay. This hybrid is equivalent to cI<sub>N</sub>-Inv<sub>pstb</sub>478, in which the homologous domains of the C-terminus of invasin<sub>pstb</sub> were fused to  $cI_N$ . In addition, an  $inv_{pstb}$  fragment carrying codons 575 to 694, which encompasses D2 (see Fig. 2 [7]), was inserted between cI<sub>N</sub> and the *inv<sub>ent</sub>* coding segment, to mimic invas $in_{pstb}$ . The chimeric  $cI_N$ -Inv<sub>ent</sub> proteins were readily detected on SDS-polyacrylamide gels of cell extracts (Fig. 3) although the steady-state level of  $cI_N$ -Inv<sub>pstb</sub>575-694-Inv<sub>ent</sub> was somewhat lower than that of the other chimerae (Fig. 3, lane 7).

The ability of the  $cI_N$ -Inv fusion proteins to repress transcription was tested, based on the property that only multi-

meric cI is able to form a repressor of the highest activity levels (13). As expected, a wild-type  $\lambda$  repressor, a cI\_N-GCN4 chimera containing a leucine zipper dimerization domain, cI\_N-Inv\_{pstb}478, and cI\_N-Inv\_{pstb}575-694 yielded 84 to 99% repression of  $\lambda P_R$ , respectively (Fig. 2). In contrast, cells expressing the cI\_N-Inv\_{ent}380 protein yielded 52.4% repression of  $\lambda P_R$ , essentially identical to the monomeric cI\_N (41%) and cI\_N-Inv202 proteins (48%), indicating that the construction containing the C terminus of invasin\_{ent} is monomeric. The triple hybrid cI\_N-Inv\_{pstb}575-694-Inv\_{ent}380, which was less abundant in the cell compared to other fusion proteins, was still highly effective at repressing  $\lambda P_R$  (99% repression; Fig. 2).

Strains expressing chimeric proteins that result in a functional dimeric or multimeric repressor, such as cI and cI-GCN4, confer immunity against the lytic phage λKH54 (ΔcI). In contrast, clones expressing monomeric cI proteins are sensitive to the phage (13). Consistent with previous results, the cI-Inv<sub>pstb</sub> 478 and cI-Inv<sub>pstb</sub> 596-694-Inv<sub>ent</sub> 380 proteins conferred resistance to the phage, implying multimer formation, while cI<sub>N</sub>-Inv<sub>ent</sub> 380 failed to promote immunity to the phage (Fig. 2), consistent with its monomeric state.

Invasin protein lacking D2 is defective for stimulation of uptake. To determine if a lack of the D2 region has functional consequences on invasin-promoted uptake, two different internal in-frame deletion mutations removing the entire D2 region were introduced into the cloned inv<sub>pstb</sub> gene and placed into a low-copy-number vector under the control of the inducible lacPO promoter. The protein products differed in whether or not they had four Y. enterocolitica invasin residues (AVLP) that may provide a linker region in between two immunoglobulin superfamily domains [called ( $\Delta D2$ )-1 and ( $\Delta D2$ )-2; Fig. 1B]. Proteins with the expected molecular mass were detected in E. coli extracts for both invasin<sub>pstb</sub> ( $\Delta D2$ )-1 and invasin<sub>nstb</sub>( $\Delta D2$ )-2, with expression levels that were similar to the invasin<sub>pstb</sub> and invasin<sub>ent</sub> proteins using the identical vector (data not shown). Invasin derivatives from both bacterial species and the two D2 deletion derivatives of invasin<sub>pstb</sub> were tested for binding to cultured cells (see Materials and Methods). In parallel, the efficiency of bacterial uptake was determined. To do so, E. coli cells expressing the different invasin derivatives were grown under conditions in which invasin expression was either highly induced (+IPTG) or uninduced (-IPTG). Binding to HEp-2 cells by E. coli derivatives harboring these constructions was essentially identical for each derivative, with about  $4 \pm 0.5$  (mean  $\pm$  standard deviation) bacteria associated per HEp-2 cell, in either the presence or absence of invasin overexpression (Fig. 4A). In the absence of induction, the ability of  $\hat{E}$ . coli cells harboring the invasin<sub>pstb</sub> protein to enter HEp-2 cells was 10-fold higher than that of E. coli cells expressing the deletion derivatives or the invasinent homologue (Fig. 4B). This result is in agreement with our previous observations in which a deletion of D2 and D3 (Fig. 1B) in the invasin<sub>pstb</sub> protein resulted in highly defective cellular entry of Y. pseudotuberculosis (7). When the inducer IPTG was added to the growth medium, invasin function increased, as judged by the amount of bacterial adhesion and uptake by HEp-2 cells (Fig. 4). The difference in uptake efficiencies promoted by invasin<sub>pstb</sub> relative to either invasin<sub>ent</sub> or the deletion derivatives was greatly reduced by induction. Under these conditions, uptake was 30 to 50% the level obtained with full-length invasin<sub>pstb</sub> (Fig. 4B). Thus, high production levels can compensate for the lower uptake efficiencies of the Inv<sub>ent</sub> and deletion proteins.

As uptake and adhesion strongly depended on the amount of protein expressed by the bacterium, differences in the uptake efficiencies of bacteria harboring the  $inv_{ent}$  and  $inv_{pstb}$ 

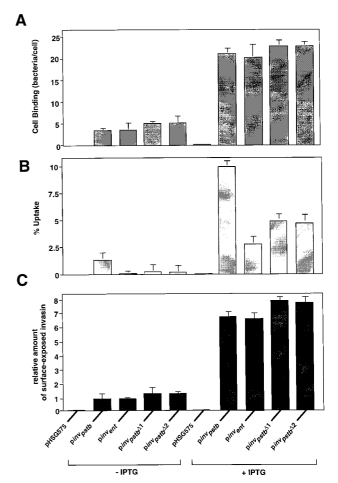


FIG. 4. Cell binding, entry efficiency, and surface exposure of different invasin alleles expressed by recombinant E. coli cells. E. coli XL1blue cells harboring plasmids pHSG575 (vector), pPD207 (inv<sub>pstb</sub>), pPD231 (inv<sub>ent</sub>), pPD254  $(inv_{pstb}\Delta D2$ -1), and pPD255  $(inv_{pstb}\Delta D2$ -2) were grown in M9 media with 0.4% glucose to an optical density at 600 nm of 0.7. Subsequently, the cultures were divided and grown for an additional 2 h in the presence or absence of 1 mM IPTG. An amount of 107 bacteria was used to infect a HEp-2 cell monolayer of about  $10^5$  cells to monitor their cell binding at  $20^{\circ}\mathrm{C}$  and to determine invasion efficiency at 37°C (see Materials and Methods). The total number of cell-associated bacteria was visualized and counted microscopically (A), and the percentage internalized is shown (B). To compare the surface concentration of the invasin alleles on the bacterial cell surface,  $2\cdot 10^7$  bacteria were used for surface ELISAs probing with either monoclonal antibody mAb3A2 against invasin<sub>pstb</sub> or with a polyclonal antiserum against invasin<sub>ent</sub> (kindly provided by V. Miller). Identical ELISAs performed in parallel with plastic wells coated with purified MBP-Inv<sub>ent</sub> and MBP-Inv<sub>pstb</sub> proteins were used to standardize the color reaction and determine the number of invasin molecules in the bacterial outer membrane. The values shown are expressed relative to the amounts of surfaceexposed invasin of bacteria XL1blue pPD207 (inv<sub>pstb</sub><sup>+</sup>) without IPTG, defined as 1.0 (C).

genes could result from either differing protein expression levels or export efficiencies. To examine whether the different derivatives were exported equally to the outer membrane, a surface ELISA technique was performed to determine the number of surface-exposed invasin molecules in the bacterial outer membrane. *E. coli* cells harboring the invasin derivatives were grown under the same conditions used for the cell binding and invasion assays, and the amount of invasin on the bacterial cell surface was quantified (see Materials and Methods). Similar numbers of the Inv<sub>ent</sub> and the Inv<sub>pstb</sub> proteins were found in the outer membranes of *E. coli* cells, although the amount of surface-localized protein was approximately 10% higher for

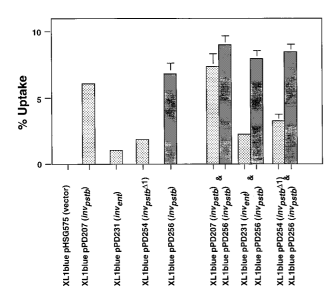


FIG. 5. Competitive uptake assays using two recombinant *E. coli* strains expressing either wild-type invasin<sub>pstb</sub> or another invasin derivative. *E. coli* XL1blue cells harboring plasmids pHSG575 (vector), pPD207 (*inv*<sub>pstb</sub>), pPD231 (*inv*<sub>ent</sub>), pPD254 (*inv*<sub>pstb</sub>ΔD2-1), and pPD256 (*inv*<sub>pstb</sub>) were grown in M9 media with 0.4% glucose to an optical density at 600 nm of 0.7. Subsequently, the cultures were grown for additional 2 h under inducing conditions with 2 mM IPTG. For the competition experiment, equal amounts of XL1blue pPD256 and XL1blue cells harboring either pPD207, pPD254, or pPD231 were mixed and 10<sup>7</sup> bacteria were used to infect a HEp-2 cell monolayer of about 10<sup>5</sup> cells at 37°C (for details see Materials and Methods). The percentage of single or mixed bacteria internalized is shown. Error bars represent the standard deviations of the means of four groups of 50 cells counted in various regions of the stained specimens, and data presented are one of three typical independent experiments.

the deletion derivatives (Fig. 4C). This relationship between derivatives was maintained in either the presence or absence of the inducer IPTG (Fig. 4C). Hence, the differences in the uptake efficiencies of the invasin homologs must be explained by the distinct properties of these proteins and not by differences in expression or presentation of proteins on the bacterial cell surface.

Inefficient competition by overexpressed invasin derivatives lacking D2. To examine if, under conditions of overproduction, bacteria harboring inv<sub>pstb</sub> are indeed more efficient at promoting uptake than bacteria harboring invent, competitive uptake experiments were performed. An ampicillin resistance cassette was inserted into the chloramphenicol resistance gene of pPD207 (inv<sub>pstb</sub><sup>+</sup>) to distinguish between E. coli strains expressing the invasin<sub>pstb</sub> wild type or the deleted invasin alleles. Subsequently HEp-2 cells were coinfected with either equal amounts of recombinant E. coli XL1blue pPD256 (inv<sub>pstb</sub><sup>+</sup>) and XL1blue pPD231  $(inv_{ent}^+)$  or XL1blue pPD254  $(inv_{pstb}^-)$   $(\Delta D2)$ -1) after growing the strains under inducing conditions (+IPTG). The uptake efficiencies of the deletion derivatives were less than 40% that of the wild type (Fig. 5). This parallels the results of experiments using HEp-2 cells challenged with single bacterial strains. Therefore, even under conditions that maximize expression of the deletion derivatives, the proteins lacking D2 were less active than invasin<sub>pstb</sub>.

## DISCUSSION

This study demonstrated that in contrast to *Y. pseudotuberculosis*, the invasin protein of *Y. enterocolitica* which lacks the self-association domain D2 was not capable of promoting homotypic interaction. This result was also supported by in vivo

cross-linking experiments, in which efficient cross-linking of high-molecular-weight complexes of invasin could only be seen with invasin<sub>pstb</sub> but not with derivatives lacking D2 (data not shown). To examine if the lowered efficiency of invasin<sub>ent</sub> multimer formation affected its function, we compared cell attachment and uptake mediated by the invasins from these two *Yersinia* species or to in-frame deletions of invasin<sub>pstb</sub> engineered to resemble the natural deletion in invasin<sub>ent</sub>. In all cases, *E. coli* cells carrying the invasin<sub>pstb</sub> clone were significantly more proficient at uptake than *E. coli* expressing the deletion derivatives, although high-level expression of derivatives lacking D2 significantly overcame this difference. Therefore, we conclude that D2 is a critical determinant for promoting uptake under conditions in which invasin concentration is limiting.

Functional differences between the *Y. pseudotuberculosis* and *Y. enterocolitica* invasin have been observed previously (28, 45). Miller and coworkers found that the uptake efficiency of  $inv_{em}$ -harboring strains was 6- to 60-fold lower than that of  $inv_{pstb}$  strains, depending on the cell line being analyzed. The results obtained here are also consistent with a previous analysis of D2 (7), in which far more internalization was observed with beads coated by an invasin derivative containing D1 to D5 than by D4 and D5 alone. Thus, lack of multivalency seems to strongly correlate with a decrease in the ability to be internalized by mammalian cells. This result emphasizes the importance of invasin-invasin interaction for efficient internalization.

We propose that interaction of homomultimeric invasin with multiple integrin receptors establishes intimate adherence and receptor clustering, providing a signal for internalization. It has been shown that the interaction of extracellular matrix proteins with  $\beta_1$  integrins leads to the association of multiple signaling proteins and cytoskeletal elements at the site of binding to substrate (43). The invasin-bound integrin complex is still uncharacterized, although it appears to consist of several tyrosine-phosphorylated proteins, such as Paxillin, Cas<sup>130</sup>, and FAK (1; P. Dersch, unpublished results). As monomeric and multimeric forms of invasin lead to different cellular responses (7), there may be differences in the signaling molecules associated with the integrin after adherence with each form. Furthermore, the concentrations of signaling proteins recruited to the cytoplasmic domain of  $\beta_1$  integrins may depend on the multimerization state of the protein, affecting relative uptake efficiencies.

The relative efficiency of bacterial uptake mediated by different invasin alleles strongly depends on the amount of surface-localized invasin molecules (see Fig. 4C). Under noninducing conditions, under which a small number of invasin<sub>pstb</sub> molecules was expressed on the surface of *E. coli* cells, uptake was approximately 6- to 10-fold higher than uptake of bacteria having invasin derivatives lacking D2. High expression levels, however, greatly suppressed this difference in uptake efficiency, allowing compensation for the absence of D2. This finding agrees well with previous observations using invasin-coated latex beads, in which invasin<sub>pstb</sub> having only D4 and D5 (Fig. 1A) required more than 10,000 molecules per bead to promote efficient uptake (7). This coating concentration is approximately 10 times higher than that necessary to promote uptake by D1 to D5.

It is unclear what pressures resulted in *Y. enterocolitica* and *Y. pseudotuberculosis* having invasin molecules with different numbers of immunoglobulin superfamily repeat modules. In addition to invasin, differences between adhesion proteins of *Y. enterocolitica* and *Y. pseudotuberculosis* have been observed. *Y. enterocolitica* Ail was first identified as a protein that mediates cell adhesion and low-level uptake (29). In contrast, the *Y.* 

pseudotuberculosis Ail protein promotes neither cell binding nor cell entry (44). Furthermore, the plasmid-encoded adhesion factor YadA of Y. enterocolitica seems to be more efficient in tissue culture uptake experiments than its Y. pseudotuberculosis homologue and has been demonstrated to play a more critical role in Y. enterocolitica virulence than that observed with Y. pseudotuberculosis (34). Thus, it seems very likely that although the Y. enterocolitica invasin is less competent at promoting uptake than invasin<sub>pstb</sub>, this has been compensated for by features in proteins such as Ail and YadA that allow them to be more effective at cellular interaction than their Y. pseudotuberculosis counterparts. It is likely that these differences have consequences on the relative courses of infection taken by these two species. Yersinia species have fairly broad host ranges and a particular adhesion factor may be more effective in one animal host than in another. A second explanation for differences in activities of similar proteins is that the two bacterial species may use the identical adhesive factors for different purposes. Perhaps even the tissue site of action of a particular adhesin differs between the two bacterial species.

Slight alterations in invasin function may explain differences in the colonization pattern of the enteropathogenic Yersinia species and may also be responsible for their distinct pathogenicities. Both Y. enterocolitica and Y. pseudotuberculosis translocate through the intestinal epithelial layer via M cells overlaying the Peyer's patches. Subsequent colonization of the underlying lymphatic tissue seems to be significantly different, and may be altered by relative invasin function. It has been observed that Y. enterocolitica cells form abscesses associated with large numbers of bacteria in the mouse Peyer's patch, while infections with Y. pseudotuberculosis yield much smaller numbers of bacteria with no abscesses in this site (Ingo Authenrieth, personal communication; P. Barnes and R. Isberg, unpublished results). It is possible that the increased activity of invasin<sub>pstb</sub> may allow more efficient bypass of antiphagocytosis promoted by YopH and YopE, with the consequence of reducing the proliferation of Y. pseudotuberculosis within lymph nodes. Future analysis of animal infection models is needed to fully evaluate the consequences of differences in the properties of adhesive protein encoded by enteropathogenic Yersinia species.

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